

# Machine Learning at Scale

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## ABSTRACT

It takes skill to build a meaningful predictive model even with the abundance of implementations of modern machine learning algorithms and readily available computing resources. Building a model becomes challenging if hundreds of terabytes of data need to be processed to produce the training data set. In a digital advertising technology setting, we are faced with the need to build thousands of such models that predict user behavior and power advertising campaigns in a 24/7 chaotic real-time production environment. As data scientists, we also have to convince other internal departments critical to implementation success, our management, and our customers that our machine learning system works. In this paper, we present the details of the design and implementation of an automated, robust machine learning platform that impacts billions of advertising impressions monthly. This platform enables us to continuously optimize thousands of campaigns over hundreds of millions of users, on multiple continents, against varying performance objectives.

## Categories and Subject Descriptors

I.5.2 [Computing Methodologies]: Pattern Recognition

## General Terms

Design, Reliability, Measurement

## Keywords

machine learning, predictive modeling, scalability, computational advertising

## 1. INTRODUCTION

Demand for digital advertising from consumer brands has been growing rapidly in the past few years because of a recent shift to more and more content becoming accessible to people on digital devices. Digital advertising spans display (banner), video and rich media ads that are shown to users

while they are browsing the Internet, as well as ads shown in mobile apps and on addressable TV sets. In contrast with traditional TV advertising, which is generally reaching households within the intended demographics and geographic location, digital advertising offers the advantage of targeting specific users and measuring their responses. This is frequently implemented by creating pools of browser cookies that anonymously represent people within the target audience for an advertising campaign. When a web page with space allocated for an ad loads in the web browser, the browser generally sends a request for an ad along with the user's browser cookie to an ad server. Part of the decision by the ad server of which ad to send back to the browser depends on whether the cookie is in a targeted pool.

A common way of generating pools of users is collecting cookies of people who expressed interest in a given product or service by visiting the advertiser's web site (known existing or potential customers). However, advertisers are also interested in finding other people for whom their ad may be relevant (unknown potential customers). The latter is frequently addressed by building a model predicting which cookies belong to people who are in the target audience for a given campaign [7, 2, 3, 1].

The target audience for a given advertising campaign can be defined in different ways, largely depending on the campaign goals. In general, these goals are of two types: reach and performance. Campaigns with reach goals are targeting a certain audience segment, e.g., people likely to watch a given TV show, or likely to be interested in sports. Campaigns with performance goals, on the other hand, are targeting users who are likely to complete a certain action in the future, e.g., click on the ad or purchase a product online after viewing the ad.

At Collective we are faced with the challenges of building, testing, maintaining and keeping track of thousands of such models that enable daily delivery of tens of millions of ad impressions for hundreds of advertising campaigns. The data involved occupies hundreds of terabytes of storage, it is time-dependent and comes from multiple and evolving data sources in different formats. The campaigns run continuously (24/7) on our platform and are routinely added, removed or modified. Finally, the resulting user pools must be kept up-to-date in the real-time systems that deliver ads. This paper describes the system currently deployed in production at Collective that enables building and maintaining thousands of predictive models at a time, and making hundreds of billions of predictions on a daily basis.

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## 2. MODEL SETUP

In this section, we define the goals of our modeling platform, discuss the time structure of the data used for model building, outline the features used to make predictions and describe our choice of modeling methodology.

### 2.1 Goals of Modeling

The modeling needs of campaigns with performance and reach goals are very similar, however, for simplicity we will only describe models for campaigns with performance goals. When an ad is sent by the ad server to the user’s browser (an event referred to as an “impression”) there is a chain of other events that follows. First of all, the user may or may not see the ad, depending on whether the ad is above or below the fold of the web page and is “viewable”. A standard for measuring ad viewability has emerged recently to mean “50% or more of the ad was visible on the screen for 1 second or longer” [6]. If the ad is viewable, the user may interact with the ad in some way. For example, the user may bring the mouse cursor to the ad, or, in case of video ads, view it to completion or skip it. Then, the user may click on the ad and follow the link to the advertiser’s web page. Finally, the user may “convert”, which can mean buying a product, signing up for an online newsletter, requesting information for a product, etc. Note that conversions may happen regardless of whether or not the user interacted with or clicked on the ad.

In the chain of events, from the ad being available in the browser, to view, to interaction, to click, to conversion, the chances of the desired event happening go down, while the value of the event to the advertiser goes up. Every campaign with performance goals has one or more goals within this chain of events. As a result, we frequently have to build more than one type of model per campaign. In addition, when the target events are too rare to build a meaningful model, which may happen at the very beginning of a campaign with no prior history, the system automatically falls back to a previous event type in the chain that has a higher frequency of occurrence. For example, a model for predicting clicks would fall back to a model predicting user interactions with an ad.

In order to create a unified modeling process across all event types, we take advantage of the fact that in all cases the response variable in the model can be treated as a binary variable that identifies whether or not a given event happened within a predefined time interval  $T$  after the ad was served. Thus, for each model we are predicting the probability of a given user  $U$  to complete a specific action  $A$  within time  $T$  (“look-forward window”) after being served an ad. This formulation can be readily extended to the response variable representing counts of events, and the model predicting the expected number of user actions.

### 2.2 Time Structure of Data

Because our models are predicting events occurring in the future, the response variable has to be measured later in time relative to the state of the predictor variables. Figure 1 shows a diagram of the time structure of the predictor and response variables. The users are observed over a period of time, at the end of which predictions are made and the users are assigned to the target user pools. There is a short delay arising from the time taken to deliver the data to real-time systems. Once the user pools are available for targeting on

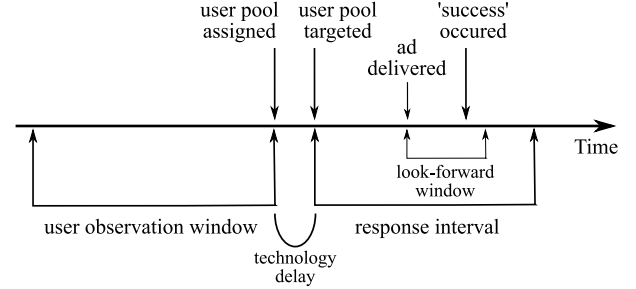


Figure 1: Time relationship between predictor and response variables.

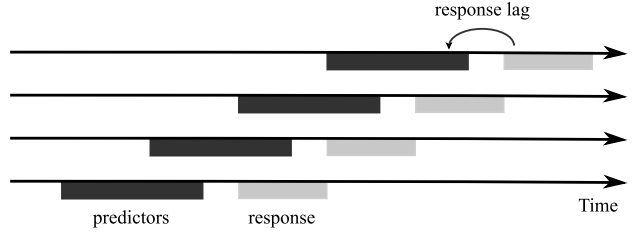


Figure 2: Time structure of the training data.

the ad server, the response interval begins. The response interval accounts for all occurrences of ad delivery and the events that are considered a success for the given campaign within a look-forward window after an ad was delivered.

The same time structure of the data is reflected in the training data set, as shown in Figure 2. We use the information that we knew about a user at the end of the observation window as predictor variables, while the response variable indicates whether or not the user has performed the desired action given that we delivered an ad to this user. We construct a sequence of non-overlapping response intervals of equal length, except that the latest response interval may be incomplete in order for the model to take advantage of the most recent available data. Note, however, that the look-forward interval for capturing the response remains consistent to avoid biasing the outcome rate in the most recent interval.

Typically, the user observation period is four weeks, the response interval is one week, and we use a set of 8 such intervals to construct the training data set. The number of response intervals is a tradeoff between the amount of historical data we have to store, its relevance to the current user’s behavior, and the number of positive events available to the model. The data in the response intervals is exponentially weighted so that more recent data contributes more to the model than the older data. In addition, in cases where the number of success events is very large, only events from the most recent intervals are included in the training set.

### 2.3 Predictor Variables

The data that is meaningful as features for predicting user behavior comes primarily from Collective’s delivery of billions of ads on behalf of our clients every month. Such features may include web site visitation history, geographic in-

Table 1: Strengths and limitations of glmnet

Component	Strengths	Limitations
Logistic regression model	<ul style="list-style-type: none"> <li>· Produces probabilities*</li> <li>· Speed of predictions*</li> <li>· Interpretability of coefficients</li> </ul>	<ul style="list-style-type: none"> <li>· Nonlinearity must be specified*</li> <li>· Interactions must be specified*</li> <li>· Limited missing value support</li> </ul>
Elastic net regularization	<ul style="list-style-type: none"> <li>· Reduces risk of overfitting*</li> <li>· Parsimonious with <math>\alpha</math> near 1</li> </ul>	<ul style="list-style-type: none"> <li>· Limits statistical inference</li> <li>· Requires choosing <math>\alpha</math></li> </ul>
Coordinate gradient descent	<ul style="list-style-type: none"> <li>· <math>\lambda</math> search path for “free”*</li> <li>· Sparse data support*</li> </ul>	
R package implementation	<ul style="list-style-type: none"> <li>· Cross-validation for <math>\lambda</math> search</li> <li>· Ease of use of R interface</li> <li>· Speed of model building*</li> </ul>	<ul style="list-style-type: none"> <li>· Not parallelized except for cross-validation</li> </ul>

\* These strengths (and limitations) are most significant, and are explained in the text.

formation derived from the IP address, device and browser information inferred from the user-agent string within the browser HTTP request, and other data available in the process of ad delivery. This data is both high-dimensional and sparse. One can think of this data set as an  $n \times p$  matrix where each of the  $n$  rows represents a user, and each of the  $p$  columns represent a feature of the users. In the following, we’ll interchangeably refer to this matrix as “model matrix” or “feature matrix”. Since the user features are independent of the campaigns or the campaign goals, we construct a single global model matrix for all models, providing the flexibility, however, for any particular model to add or remove features from the global model matrix. We address the challenges and our implementation of maintenance and assembly of the model matrix in detail in Section 3.

## 2.4 Choice of Modeling Methodology

Collective’s audience modeling platform is primarily concerned with predicting the probability of an event occurring for a user in the future given a vector of data known about that user at the time of the prediction. The system described in this paper uses elastic nets to make these predictions, which are regularized generalized linear models. To train these models, we use the R `glmnet` package [4], which is an implementation of the coordinate descent algorithm described in more detail in [5]. In this section, we will describe elastic nets at a high level, and then explore the strengths and limitations of these algorithms for our application.

Let our response variable be denoted by  $Y = \{0, 1\}$ , where 1 indicates that the event occurred in the future, and 0 indicates that the event did not occur. Further, let  $X$  be an  $n \times p$  matrix of data where each row corresponds to the predictors for a user. Then the elastic net solves for a set of coefficients for the logistic regression model of the form

$$\log \left( \frac{\Pr(Y = 1|x)}{\Pr(Y = 0|x)} \right) = \beta_0 + x^T \beta.$$

That is, the log of the odds of an event occurring is expressed as a linear combination of the features in  $X$  and a coefficient vector  $\beta$ . Traditionally the coefficients for a generalized linear model are determined by identifying the vector  $\beta$  that maximizes the log likelihood equation

$$l(\beta_0, \beta) = \frac{1}{N} \sum_{i=1}^N y_i \cdot (\beta_0 + x_i^T \beta) - \log(1 + e^{\beta_0 + x_i^T \beta}).$$

However, when  $p$  is large (many predictors) in proportion

to  $n$  (number of observations), maximizing the likelihood directly risks significant overfitting and poor out of sample model performance. Thus, the elastic net seeks coefficients that minimize

$$\min_{(\beta_0, \beta) \in \mathbb{R}^{p+1}} l(\beta_0, \beta) + \lambda P_\alpha(\beta),$$

where  $\lambda$  is a regularization parameter, and  $P_\alpha(\beta)$  is a coefficient penalty term defined as

$$P_\alpha(\beta) = (1 - \alpha) \frac{1}{2} \|\beta\|_{\ell_2}^2 + \alpha \|\beta\|_{\ell_1}$$

for a parameter  $\alpha$  satisfying  $0 \leq \alpha \leq 1$ . The choice of  $\alpha = 0$  corresponds to using a ridge regression penalty function, and the choice of  $\alpha = 1$  corresponds to using a lasso penalty function. Any intermediate value of  $\alpha$  is a compromise between the two. In practice, we found a value of  $\alpha = 0.1$  to provide optimal out-of-sample performance for our models with limited variation across datasets or time.

We chose the R `glmnet` implementation of coordinate gradient descent for elastic nets for a variety of reasons. First, elastic nets perform well on out of sample data, even when compared to much more sophisticated methods like random forests and gradient boosting machines. In part, this is due to the high dimensionality, sparseness and noisiness of the data that we are using. Second, elastic nets produce models which are easy to predict quickly at massive scale, due to their relative simplicity. Third, the `glmnet` implementation is both easy to use, and is incredibly fast and memory efficient. Table 1 lays out the specific strengths and limitations of this choice, and we explain the most significant strengths and limitations below.

**Logistic regression model (Strength): Produces probabilities.** The logistic regression model produces probabilities for success outcomes directly. These probabilities can be helpful in interpreting the model predictions directly, but more importantly can be used in downstream applications. For example, groups of users can be created that exceed probability thresholds, and the probabilities can be used directly in ad serving or real time bidding applications.

**Logistic regression model (Strength): Speed of predictions.** Every day our system must score every user for every model still in production. Given that we have  $\sim 200$  million users, and  $\sim 1,000$  models in production at a given time, that means we are making over 200 billion predictions every day. The required computation for a given user and model is a sum-product of the coefficients and the user fea-

tures, which can be implemented very easily and performed incredibly quickly. These 200 billion predictions are done in under an hour in our current implementation.

**Logistic regression model (Limitation): Nonlinearity must be specified.** Because logistic regression assumes linearity in the additive terms of the log-odds, any more complex nonlinear relationships between predictors and the log-odds must be parameterized in the model matrix in advance. We do so by either binning continuous features, or by using nonlinear transformations such as polynomials or splines. In general, the majority of our predictor data is categorical, and so this is not a material concern for our application.

**Logistic regression model (Limitation): Interactions must be specified.** Because logistic regression assumes additivity of terms in the log-odds, any interactions effects must be specified in the model matrix in advance. With a large number of features, this quickly becomes intractable in both processing time and memory storage, so interactions must be added sparingly.

For this application, we believe that interaction terms are less important than they might be for other applications. First, we are only capturing data related to users and their past behavior. This means all of our data is directly related to a single entity (the user), and so complex interactions between multiple entities are not present. Second, many of our features are sparsely populated, and so interactions of those sparse features are themselves highly sparse, and necessarily less likely to influence the prediction. Finally, we have a very large number of features, drawn from many disparate sets of data. We believe that useful interaction effects amongst features become less useful as additional features are added that can explain those interactions directly.

**Elastic net regularization (Strength): Reduced risk of overfitting.** Regularization is a crucial feature of elastic nets. By penalizing coefficient sizes, we ensure that the models are much more likely to generalize to new data, and thus perform well in production. This enables us to more freely add large sets of sparse features without being concerned that small sample size models will overfit them. We also reduce the need to use variable selection techniques as a part of our daily model building processes, which can be computationally intensive.

**Coordinate gradient descent (Strength):  $\lambda$  search path for “free”** In a production machine learning system, one key consideration is the need to perform grid searches for algorithm meta parameters, which can be computationally costly. The choice of  $\lambda$  in elastic nets has a tremendous impact on the resulting model, with  $\lambda = 0$  producing the (overfitting) full GLM fit, and  $\lambda = \infty$  producing the (underfitting) constant model. With a wide variety of dataset sizes and outcomes to predict, we find that in practice optimal (maximum cross-validated AUC)  $\lambda$  values regularly vary from 0.0001 to 0.1. One significant advantage of the coordinate gradient descent algorithm is that it iteratively solves for the optimal coefficients for  $\lambda$  along a  $\lambda$  search path (from largest to smallest), producing the grid search for this parameter automatically, often faster than if the  $\lambda$  search was not done at all.

**Coordinate gradient descent (Strength): Sparse data support.** A further advantage of the coordinate gradient descent algorithm is its support for sparse matrices. On average, entries in our user matrices are non-zero less

than 10% of the time, and so sparse matrix support reduces the required amount of memory by a factor of 10. This allows us to use larger matrices that produce models with higher predictive performance.

**R package (Strength): Speed of model building.** The R `glmnet` package is a wrapper for a Fortran function which implements the coordinate gradient descent algorithm. This code is highly optimized for speed and is generally stable in our production system. We build tens of thousands of models every week, and so this is a significant computational cost improvement over traditional regularized glm implementations.

**R package (Limitation): Not parallelized.** The most significant weakness of this package is that the code is not parallelized. A small amount of parallelization can be achieved by conducting the cross-validation model builds in parallel, but in practice we find that we are building many such models, so parallelization is assumed at the system level across thousands of models run simultaneously.

### 3. FEATURE MATRIX

Preparing the data for use as predictors in modeling has its own challenges. First, when the data is captured, it is generally in the format of the source system and needs to be transformed into data structures that are compatible with the modeling algorithms. Second, dimensionality of the data needs to be reduced to address both the extreme sparsity of the data and the scaling of the modeling algorithms with the number of dimensions. Third, the time structure of the data requires that the features defined at one point in time could be used for constructing the data set at another point in time. Finally, the features should be defined, added, and removed without having to modify the source code. This section describes Collective’s implementation of the data assembly process that constructs the features used as predictor variables in modeling, as well as approaches we take to selecting the feature sets.

#### 3.1 Feature Types

For our application, the algorithm of choice for building predictive models is `glmnet`, as outlined in the previous section. This choice imposes certain requirements on the structure of the predictor variables. We implemented four generic types of transformations that convert the features from the format in which they are available for modeling to final predictor variables that depend on the specific data at the time when these variables are defined.

Continuous features may need to be transformed to binary format using binning to account for nonlinearity. In general, binning attempts to break a set of ordered values into evenly distributed groups, such that each group contains approximately the same number of values from the sample. In practice, one has to account for common cases when a disproportionate number of values are the same, or when the distribution of values is discrete and heavily skewed. Standard implementations of binning, such as computing quantiles, don’t perform consistently in such cases. An additional consideration is that in the context of modeling it is impractical to create a bin with very few data points because that leads to a feature that is extremely sparse and is unlikely to be valuable. We implemented a robust method for binning that performs well across a wide variety of distributions and edge cases. This method is minimizing the least

mean square deviation of the resulting number of points in each bin from an ideal split, where each bin has the same number of points, using a combination of quantiles and the operations of splitting and merging the bins. The optimization is constrained by the desired number of bins and by the minimum number of points in a single bin.

Categorical features must be transformed to binary format by creating a binary variable for each categorical value. However, high-cardinality features, such as web pages where ads were delivered, would then translate into millions of predictor variables, the vast majority of which are extremely sparse. To alleviate this issue, we apply two approaches. The first is to group related categorical values. For example, web pages could be grouped by Internet domain, or by the category of the page contents. The second is to limit the categorical values that become predictor variables in the feature matrix to only the most common ones in some sense.

In our implementation, a categorical variable can be limited by one of two methods: top coverage and minimum support. The first method, top coverage, is selecting categorical values by computing the count of distinct users for each value, sorting the values in descending order by the count of users, and choosing the top values from the resulting list such that the sum of the distinct user counts over these values covers  $c$  percent of all users, for example, selecting top geographic locations covering 99% of users. This works best with features that only allow one value per user. The minimum support method is selecting categorical values such that at least  $c$  percent of users have this value, for example, web sites that account for at least  $c$  percent of traffic. This restriction is most amenable to features with more than one categorical value per user.

To summarize, the first three transformations of the predictor variables are top coverage and minimum support for categorical variables, and binning for continuous variables. The fourth and final transformation is a trivial one of identity, where the predictor variables are taken without change, for example, when a feature is binary to begin with.

## 3.2 Dimensionality Reduction

In a typical month, Collective’s ad delivery systems encounter billions of unique browser cookies across millions of online content items. Producing a feature matrix for every user (cookie) and every piece of information about a user as an  $n \times p$  matrix, where  $n$  is the number of users and  $p$  is the number of predictor variables, is impractical both from the data processing standpoint and because the resulting matrix would only have about 1 in 100,000 non-zero elements. In our modeling system we reduce dimensionality in both  $n$  and  $p$  to arrive at a few hundred million relevant users and between one and two thousand predictor variables, with data sparsity of about 1 in 10. The final feature matrix  $X$  is stored in the sparse representation in a database table, where each row contains the user id  $i$ , the feature matrix column id  $j$ , and the value  $X_{ij}$ . This dramatically reduces the data storage and processing requirements, while representing the feature matrix in the format native to `glmnet`.

Reduction of the number of predictor variables is achieved primarily by first selecting the features that become part of the data set, and then limiting the number of resulting columns in the feature matrix with the transformations described above. Reduction of the number of users for which we compute the features arises from the fact that for a large

proportion of users we only deliver a single ad impression (they are either blocking or deleting cookies). We therefore limit the user universe to those relevant users that have a reasonable chance of being encountered in the next time period. The relevant user definition is evolving and is outside the scope of this paper. It is a balance between potential reach (how many users could be targeted), actual reach (how many users of those that are targeted will be seen on any given day), and the requirements for storage and processing power. For example, one could consider users with ad impressions at least  $\Delta t$  apart and vary the time interval  $\Delta t$  to arrive at an acceptable user universe.

## 3.3 Time-Dependent Feature Definitions

While the data sources that provide the features used for building predictive models change infrequently, the specific definitions of predictor variables in the feature matrix depend on the actual data available at the time when the models are built, because of the need for binning the values of continuous variables and for limiting the number of values for categorical variables as described above. If we chose to continuously update the feature matrix definitions, we would have to rebuild every model in our system in order to make predictions using the most current definitions. However, we found that once a sufficient training data sample is available, retraining the models doesn’t add much to the model accuracy, while consuming a lot of resources.

In our modeling platform, we chose to use the feature definitions computed at the end of the most recent user observation period, i.e., we update them every  $r$  days, where  $r$  is the length of the response interval, typically, once a week. We rebuild all models at that time as well, and make predictions using these models every day. The feature definitions comprise a set of source features included in the feature matrix, the type of each feature, and the metadata describing the feature transformation into the final predictor variables. The feature transformation metadata includes the bin boundaries and the categorical value sets, along with the assignment of each of the predictor variables to a specific column of the feature matrix. The definitions computed at a given time are used for generating the feature matrix for all users until the next update, as well as for generating the feature matrix for each of the past intervals to assemble the training sets for the modeling. This reprocessing of past intervals’ data is computationally demanding but necessary to ensure all training data is consistent.

## 3.4 Feature Selection

The choice of `glmnet` as a modeling algorithm means that feature selection is not of immediate importance in ensuring a given model generalizes well, as regularization ensures that coefficients for features unrelated to the response are close to zero. However, we still have to choose from amongst hundreds of thousands of potential features a subset to include in the modeling platform. This choice requires insight into which of the features have the greatest impact upon model performance. Further, such insight can be invaluable in steering feature engineering decisions and data collection decisions in the broader technology platform over time.

Absent regularization, ANOVA tests can be used to compute  $p$ -values for terms included in a generalized linear model. Assuming the system that generated the data conforms to the assumptions of the model (rare in our experiences), these

p-values can be a reliable way of identifying which features are most significant. For lasso models, it is possible to compute the covariance test statistic [9]. However, this is a recent development that has not been generalized to elastic nets.

Instead, we employ two different approaches for estimating the importance of features. The first is commonly referred to as a dropterm, wherein we group sets of related features (e.g., all geographic features) and compare the predictive performance of a model without those features (the 'dropped' model) to one with them (the 'full' model). We perform 5-fold cross validation to both the full and dropped models and compare their area under the curve (AUC) statistics. Feature groups whose removal does not materially reduce the AUC are considered to be 'weak' feature groups for a given model. We evaluate both the average change in AUC across all models in production as well as the distribution in change in AUC as some features will be highly important for a small number of models, but not important for the majority of others.

The above approach is too computationally intensive to run continuously in production. It requires building every model five times (for cross-validation) for every feature group, which can be in the 100s depending on the level of grouping performed. So in practice this is performed periodically on a representative sample of models as significant changes to the set of features is contemplated by our data sciences team.

However, we still wish to have a measure of variable importance continuously available in our production system. This is useful both to report on insights for individual models, and also to track any changes over time in feature group performance which might indicate upstream data availability or processing issues. We have found that a simplified approach to measuring the impact of a feature group on predictions to be correlated to the more robust dropterm approach described above but far less computationally intensive. Specifically, for every group of features we set their coefficients to zero in each model, and compare the assignments made by the altered model to those made by the original model. Given that the models are regularized, this tests whether or not the coefficients are meaningfully altering the predictions, without being overly sensitive to highly correlated data (which could be the case in an unregularized GLM).

### 3.5 Assembly

The main requirements for the system that assembles the feature matrix were to (i) define features for a given set of models without modifying the source code of the system; (ii) maintain multiple sets of features for different sets of models; (iii) persist and maintain multiple feature definitions depending on the time when they were generated; and (iv) assemble the feature matrix on demand for a given subset of users, the feature definitions, and the source data as of a specific date. We separated the feature matrix generation into three stages: setting up configuration, computing feature definitions, and producing an instance of the feature matrix. The logical separation of these stages provides the required flexibility to satisfy the requirements in production, and allows for further research and exploration of the features as new data sets become available.

First, we define which data becomes part of the feature

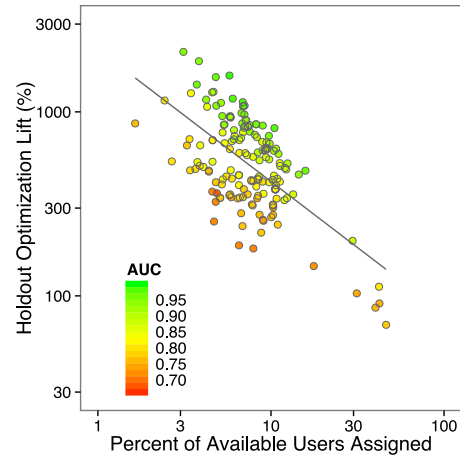


Figure 3: Evaluating model performance.

matrix. This is done through manually setting up a configuration that defines the source of every feature, the feature transformation type, parameters of the transformation, as well as other metadata that is useful in reporting and visualization. All data sources are pre-processed and assembled in a relational database (see the section on Systems Architecture below for more detail). The feature data source is most commonly a database table containing the user ids and a column with the feature values. The configuration allows for specifying custom transformations of the column values using SQL, as well as for applying filters that limit the rows included in the data set to those that satisfy the filter conditions. The parameters of the feature transformations described above include, for example, the desired number of bins and the minimum percent of values in each bin for the binning algorithm, and the percent  $c$  parameter for the top coverage and the minimum support transformations. The configurations are versioned, and we maintain separate ones for modeling user behavior in different countries.

The second stage is the generation of the specific feature matrix definitions at a particular time. These definitions rely on the data collected during the most recent user observation period and available at the end of that period. These definitions are also versioned for each configuration. We maintain a history of the feature matrix definitions so that we are able to track the changes of the definitions in time and perform system diagnostics and troubleshooting. As discussed above, the feature matrix definitions are updated every  $r$  days, where  $r$  is the length of the response interval.

The third and final stage is the generation of an instance of the feature matrix using the given feature matrix definitions and the date as of which the features data was available. We generate an instance of the feature matrix using the most recent definitions and use it for making predictions every day. In addition, during the process of assembly of the training data sets for building models, we generate the feature matrix on demand just for users that are part of the data sample used for modeling. The latter usually happens to be a small fraction of all users because of the sparsity of the positive responses and downsampling of the negative responses.

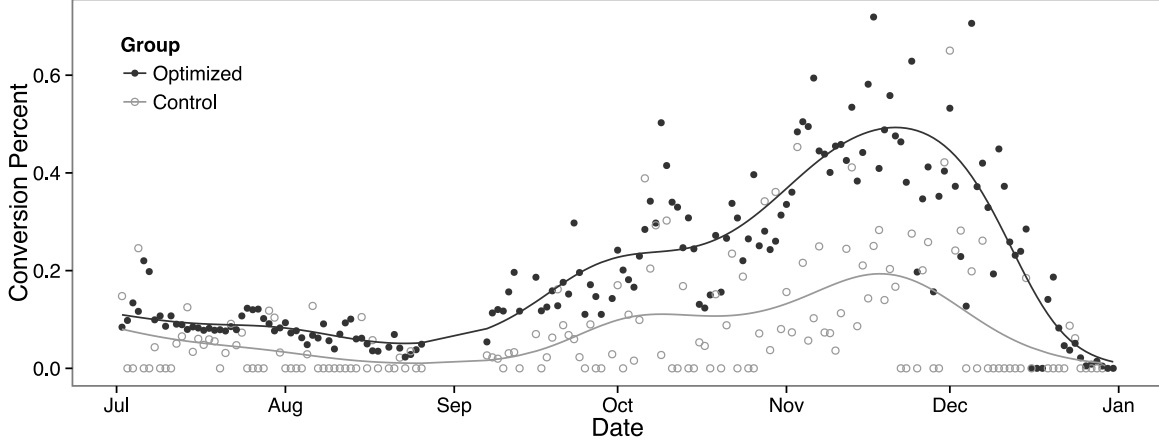


Figure 4: Sample model report of performance of optimized and control groups in production.

## 4. MODEL EVALUATION

A specific application of Collective’s audience modeling platform described in this paper is selecting a target audience for each campaign and performance goal. This is achieved by making predictions for every user in a large universe of users, sorting the predicted probabilities in descending order and assigning the  $N$  highest scoring users to the target audience. The size of the target audience  $N$  is a balance between accuracy of the predictions and the possible reach of the campaign.

To align the model evaluation methodology with the practical application at hand, we introduce the concept of “optimization lift”. We define the optimization lift  $\mathcal{L}$  as

$$\mathcal{L} = \frac{\Pr_N(\text{success}|\text{optimized})}{\Pr_N(\text{success}|\text{random})} - 1,$$

where the numerator is the probability of success (true positive rate) in the optimized set of top  $N$  users, and the denominator is the probability of success in a random set of  $N$  users.

Figure 3 illustrates the relationship between  $\mathcal{L}$  evaluated on a contemporaneous holdout set,  $N$  and the model AUC. Each point represents a model, and  $\mathcal{L}$  is on the vertical axis and the horizontal axis is the percent of available users assigned ( $N$  over the size of the relevant universe of users for each model). The color of each point is mapped to the AUC for the model. This illustration shows that the size of the target audience assigned has a dramatic effect on the performance of the model. However, given an assigned percentage of users, there remains close to an order of magnitude variation in  $\mathcal{L}$  driven by the AUC.

In our modeling platform, we take a three-level approach to evaluating the models. The first level is cross-validation to tune the model meta-parameters. In case of `glmnet` we are searching for the optimal value of  $\lambda$ . This approach has the advantage of being available at the model build time, but carries the risk of overfitting the model to the training set. The second approach is testing the model on a contemporaneous holdout set of data. The advantage of such testing is that it is available pre-deployment, but the model may still not generalize in the noisy production environment. We compute and report the AUC and the optimization lift

$\mathcal{L}$  for each model and set up alerts that send notifications for models that do not achieve a minimally required accuracy.

Finally, the third approach is embedding a random control group of users into each optimized target audience. This allows us to measure actual lift in the production environment, but it takes time and a sufficient number of impressions delivered for the campaign to achieve accuracy and statistical significance. Figure 4 shows a sample performance report, where the black dots and line represent daily performance of the optimized user group and the grey circles and line represent the daily performance of the random control group. The average lift of the optimized target audience relative to the control, computed from a sample of 58 campaigns from advertisers across different industries over a period of time in the first half of 2013 was 400%, i.e., the optimized audience performed 5 times better than random users.

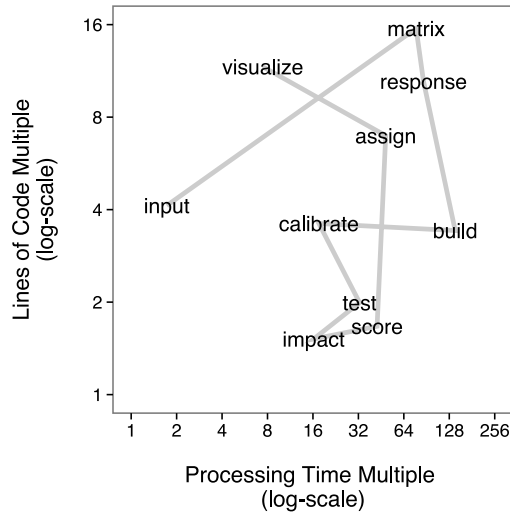
## 5. SYSTEM IMPLEMENTATION

### 5.1 Core Technologies

The architecture for a predictive modeling platform carries a number of constraints. The choice of a modeling algorithm determines which implementations are available, and that limits further choices. In our case, building the models themselves had to be done in R once we chose `glmnet`. Constructing the data sets for modeling is very intensive in data processing and requires sufficient capacity for the system. Virtually all of our data is structured and many relevant data transformations involve joins across data sets. These and other considerations led us to build the data-centric portion of our modeling platform on an IBM PureData (formerly, Netezza TwinFin) appliance. At its core is a parallel relational database with SQL interface, capable of storing and processing tables with hundreds of billions of rows. Since we needed to use both R and SQL within the system, we chose to standardize around these two languages.

### 5.2 Interaction With End Users

While the audience modeling process is implemented and maintained by the data sciences and engineering teams, the main end users of the system are the ad operations and campaign optimization teams. These teams set up the specific



**Figure 5: Lines of code involved in each step of the processing.**

inputs for each of the models, such as the campaign meta-data (e.g., campaign name), the types of the performance goals (e.g., clicks or conversions), the conversion pixel identifiers (if applicable), and the desired size of the target audience. In a separate user interface, the data sciences team sets up the global parameters of the models as well as the configurations of the feature matrix.

Upon completion of every model build, all relevant teams receive several reports detailing the results. These reports include the AUC and optimization lift for each model based on the holdout set, and flag any issues that may have occurred. For example, when a campaign has just started, there may not be enough positive events to build a meaningful model. This situation is normal, however, the optimization team receives a corresponding alert in the user interface.

As soon as the campaign has delivered enough impressions so that the optimization lift based on the embedded control group can be computed with sufficient statistical significance, a performance report similar to that in Figure 4 is produced automatically and is made available to the optimization and account services teams.

### 5.3 Modeling Process

The audience modeling process consists of 10 distinct phases, which are visualized in Figure 5 in terms of their relative processing time (indicating computational intensity) and lines of code (indicating implementation complexity). Note that the most computationally intensive tasks are highly parallelized, and so in reality are much more computationally intensive than they appear in the above chart. Each phase is briefly described below.

**Input.** The input phase is processing and structuring user input, both specific to individual models, and the meta-data controlling the entire process ( $\approx 50$  parameters)

**Matrix.** This phase assembles the feature matrix for training and prediction, and was described in detail in Section 3.5.

**Response.** The response stage assembles the response

data for all models and performs relevant sampling. Here we achieve significant savings in processing time by taking advantage of similarities in the structure of the data across model types and performing many of the data transformations on the whole data at once. There are two types of data sampling that we apply for efficiency. First, we limit the maximum number of rows in the training data on the basis of evaluating the impact of adding more data on the model accuracy relative to the additional resources for extra storage and computation. Second, in the most common case when the negative events in the response greatly outnumber the positive (assuming binary response, as described in Section 2.1) we downsample the negative events to improve speed and avoid potential numerical instability when building the model.

**Build.** The build phase involves combining the feature matrix and the response into the training data sets, distributing the modeling tasks to a cluster of computers, training the `glmnet` models, and then collecting and structuring the results. Although the `glmnet` implementation in R is not parallelized, the large number of models that we need to build at a time allows us to achieve parallelism by building a number of models simultaneously. For this purpose, we leverage our Hadoop cluster. Note that this is not a typical big data problem solved with Hadoop. Instead, we treat Hadoop worker nodes as parallel processors while taking advantage of its architecture. The modeling jobs are implemented as a mapper-only map-reduce job, where the mapper is generated by the audience modeling platform and calls R to build a single model.

**Calibrate.** This stage provides an optimization by making adjustments to the model coefficients to speed up the downstream scoring of all users for every model. Since the final result of the process for each model is a set of  $N$  highest scoring users and the model is linear, we adjust the coefficients so that the top  $N$  users have positive scores, while the rest have negative scores. This adjustment can be estimated, for example, by scoring a sample of users. As a result, there is no need to sort the complete set of scores for every model to determine the top  $N$  users. Moreover, during scoring only information about users with positive scores needs to be stored, which leads to significant savings in the amount of data written to disk.

**Test.** The test phase performs out-of-sample testing on a contemporaneous sample of data.

**Impact.** The impact phase calculates the impact of each feature group on predictions by setting the model coefficients for the features in the group to zero and comparing the resulting predictions to those made by the unmodified model. As described in Section 3.4, this serves as a proxy for the full dropterm feature selection approach.

**Score.** The scoring phase is making predictions using each model for each relevant user. Since the `glmnet` models are linear, a user's score from a given model is the sum-product of the model coefficients and the values of the feature matrix for this user. The computation of the scores for multiple models and all users can be viewed as a matrix multiplication problem: if the number of models is  $q$ , the resulting matrix of user scores  $S$ , of size  $n \times q$  is equal to  $X \times C$ , where  $X$  is the  $n \times p$  feature matrix, and  $C$  is the  $p \times q$  matrix of model coefficients. Each column of  $C$  contains the model coefficients for a single model. While  $n$  is on the order of hundreds of millions users, both  $p$  and  $q$  are on



the order of 1000. Thus, the scoring problem is equivalent to multiplying a matrix on the order of 100 million rows by 1000 columns with another matrix of size about  $1000 \times 1000$ .

Although matrix multiplication is well understood, the naive approaches did not perform well in this case. In our implementation of scoring, we take advantage of several factors. First, because the matrix of coefficients  $C$  is relatively small, one can replicate it on multiple nodes of a parallel architecture, and compute scores for distinct sets of users on each node. Second, an algorithm that takes into account the fact that both  $X$  and  $C$  are sparse can reduce the number of operations required to compute the scores by orders of magnitude. Finally, as we noted above in the description of the calibration phase, one can drastically reduce the time needed to write the results of calculations to disk by limiting the output to only relevant values.

**Assign.** The assignment phase includes performing the final assignment of users to target audiences and delivery of the results to real-time systems for targeting. At this stage we embed the control group in the target audience. We also optimize the size of the data assembled for delivery, for example, by computing and only sending the differences in the user assignment since the last model build.

**Visualize.** Each modeling process run concludes with producing a comprehensive set of visualizations for each phase of the current run, as well as time series views of prior runs. All visualizations are automated and utilize the `ggplot2` R package [10].

Not surprisingly, the tasks that require the most computational time are either incredibly data intensive (matrix and response assembly) or CPU intensive (model building). That the scoring phase is comparatively less computationally intensive is a testament to how fast `glmnet` models can be scored, and to numerous optimizations that we implemented.

The data assembly tasks (matrix and response) are also very logically intensive, as there are many decisions that must be made in those phases. However, their position is somewhat inflated by the fact that much of the code is written in SQL, a relatively verbose language (compared to `plyr` in R for example). The visualization phase is in many ways one of the most complex, as we generate thousands of visualizations for every run, of at least 50 different varieties.

## 5.4 Measurement and Monitoring

For any large scale system to be robust, it needs to implement automated testing, monitoring of correctness and performance of individual components, as well as measurement and recording of key metrics over time. It should also fail gracefully when errors and changes in data inevitably occur. An excellent overview of the design principles of large predictive modeling systems has been presented in [8]. Below we describe some of the measures implemented in our modeling platform to ensure robustness and provide for ongoing improvements.

**Integration testing.** When changes are made to such a large and complex system consisting of dozens of interdependent steps, it is difficult to anticipate all effects on different parts of the system. We have addressed this by testing the code changes on a test data set with a few representative models, and verify that the data produced and the performance of the models on the holdout set are consistent with the introduced changes.

**Timing.** For the whole system to be scalable, each component must run in within acceptable time limits. We take the development approach where we design each component to be fast but without excessive complexity, measure their execution time, and iterate on the bottlenecks. To be able to follow this approach, we measure and record the timing of every step of the process, down to individual SQL queries. Because for many tasks we are using R to run SQL queries in the database, we have implemented wrappers that can automatically record the timing of every query.

**Model performance testing** The models are tested at three levels as described in detail in Section 4.

**Monitoring key metrics.** We store a history of the most important measurements and thus make it possible to monitor the system performance and scaling over time. When key metrics change unexpectedly, we investigate and take action, as necessary.

**Error checking.** We have two key layers of error checking in the system. The first layer is system-wide, where at every phase of the multi-step process we run multiple diagnostics against the intermediate data available at that step. These tests may be as simple relational integrity checks or more complex tests based on custom logic. Whenever these system-wide checks fail it is indicative of a systemic issue that typically needs to be addressed in the code. The second layer is model specific, and captures any errors that arise due to input inconsistencies or missing or incomplete data associated with a specific model.

**Error handling.** There are multiple severity levels of errors that may occur. First, and the most severe one, is an error that causes the whole modeling process to halt. This could be an infrastructure failure, such as a network interruption or disk failure. These errors are very rare but the most disruptive, as multiple teams have to get involved to resolve the issue and restart the automated processes. Second, there are errors resulting from the required input data not being present, for example, when there is a problem with data transfer from one of the many data sources required for the data assembly. The system generally deals with this by waiting for some time for the data to arrive, as well as by sending out alerts. Finally, there are errors that occur during the modeling process. These are anticipated, diagnosed and flagged in the system and presented to end users without disrupting the flow of the overall modeling process. In addition, in cases of model specific errors, the system may fall back to another relevant model type to assign the target audience.

**Visualization** In addition to the error checking described above, we automatically generate hundreds of visualizations covering every stage of the modeling system. These visualizations help build our intuition for what a “healthy” state of the system looks like, and are thoroughly reviewed anytime a meaningful change is made to the system to look for introduced anomalies. Having these visualizations available at the individual model level, at the system component level, as well as across the whole system, proved to be critical to understanding and monitoring of the system behavior, proposing improvements, and troubleshooting.

## 6. CONCLUSIONS

Building a single accurate and scalable machine learning model to predict audience behavior for an advertiser given hundreds of terabytes of data covering hundreds of millions

of users and millions of potential features is a challenge. Architecting, implementing and supporting a system to build thousands of such models, and making certain that they run daily to ensure the proper delivery of billions of advertising impressions monthly is even more challenging. Through careful choices in data assembly, algorithm implementation and system controls and monitoring the platform described in this paper has enabled the accurate delivery of billions of advertisements in multiple countries on behalf of Collective's clients.

In this paper, we focused on predicting user actions in the context of digital advertising campaigns with performance goals. However, the same modeling platform, with minor modifications, is working for campaigns with reach goals, predicting whether a given user is likely to belong to a certain group of people. In addition, because our models produce probabilities for success outcomes, the platform has been extended to generate inputs for calculation of bid amounts for buying ad impressions at real-time bidding (RTB) ad exchanges. The described system can be further generalized to applications beyond digital advertising, in any situation where one aims to predict user behavior with multiple, possibly interdependent, outcomes. Such applications may include, for example, e-commerce and digital publishing.

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